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(54) FABRIC-SOFTENING COMPOSITIONS

We, UNILEVER LIMITED, a company organised under the laws of Great Britain, of Unilever House, Blackfriars, London E.C.4, England, do hereby declare the invention for which we pray that a patent may be granted to us and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to tertiary amine compositions useful as fabric softeners.

In recent years compositions capable of improving the softness of laundered fabrics have been sold for domestic use. Usually such fabric-softening compositions have taken the form of an aqueous solution or dispersion of a compound which confers a softening effect on fabric, and these solutions or dispersions are added to the water in which the fabric articles are rinsed after washing. A complete laundering process consists of a wash cycle in which a fabric is washed with an aqueous solution of a detergent composition, followed by removal of the bulk of the wash liquor from the fabric, and a rinse cycle, in which the fabric is rinsed with water to remove residual detergent composition, followed by removal of the rinse liquor. The rinse cycle may be repeated before the laundered fabric is dried. The need to add the softening composition during the rinse cycle can be inconvenient, particularly with automatic washing machines which would otherwise perform the complete laundering without interruption.

Attempts have therefore been made to develop a wash-cycle composition, that is, a composition to be added in the wash cycle and which is capable of softening fabrics in the presence of detergent-active compounds so that the fabric can be cleaned and softened simultaneously. This provides a difficult prob-

lem, for the active softening ingredient has to be capable of deposition in opposition to the action of the detergent whose function is to remove soil from the fabric, and in an amount sufficient to soften the fabric, but not so great that the appearance of the fabric is impaired. Furthermore, the active softening ingredient should not depress the detergent activity of the detergent-active compound in the composition, and most compounds which are known to be fabric softeners are cationic and form insoluble addition products with conventional anionic detergent-active compounds. Proposals have been made to solve this problem by using as softening agents certain betaines which undergo a charge reversal when the pH changes from the alkaline conditions of a wash liquor where they are anionic to the less alkaline conditions of a rinse liquor in which they are cationic and their deposition on to fabric is promoted. Such a betaine is soluble in the wash liquor and the small proportion of that used which is in the wash liquor residue present in the first rinse cycle has to suffice to provide the softening.

It has been proposed in German Patent 722,281 to use as textile softeners waterinsoluble primary, secondary and tertiary fatty amines in conjunction with detergents and inorganic salts, the only tertiary amine specifically mentioned being dimethyldodecylamine. British Patent 1,052,847 describes the use in the wash cycle of solid fabric softeners which are complexes of urea and primary, secondary and tertiary amines having at least one 75 straight-chain organic radical containing 12 to 18 carbon atoms, but no specific tertiary amines are mentioned.

It has now been discovered that a small group of tertiary fatty amines possess special 80

advantages as wash-cycle fabric softeners because, like the betaines referred to above and unlike dimethyldodecylamine and tertiary or other fatty amines in general, they are negatively charged under the relatively strong alkaline conditions present in a typical wash cycle and therefore do not complex with anionic detergent-active compounds, but are protonated and therefore cationic and fabric-10 substantive under the neutral or slightly alkaline conditions present in a rinse cycle. They can therefore be added to an alkaline detergent wash liquor without reducing its cleaning capacity, and as they are insoluble in the wash liquor, when this is discharged most of the amine is left behind entrapped in the fabric so that it is available for softening in the rinse: this residual amine becomes substantive to the fabric when the rinsing water is added, so that before the rinse liquor is discharged it is adsorbed on to the fabric.

The isoelectric point of such an amine lies within narrow limits, for the change in its electronic properties, or charge reversal, has to occur between the pH of the wash liquor and the pH of the first rinse liquor, which is still weakly alkaline due to residual wash liquor. The pH of a wash liquor containing anionic detergent-active compounds is usually within the range from 8.0 to 9.9, depending on the detergency builder employed. It has been found that where the isoelectric point of the amine is substantially below the pH of the wash liquor and at least 0.2 units above the pH of the rinse water the charge reversal occurs on addition of rinse water to the fabric after discharge of the wash liquor. The amines having the desired properties are tertiary amines having an isoelectric point of from 8.3 to 9.8 and the structure RR¹R²N where R is an alkyl group having from 1 to 6 carbon atoms and R1 and R2 are primary linear alkyl or alkenyl groups having from 10 to 26 carbon

The isoelectric point of any given amine can be determined by measuring the electrophoretic mobility of an aqueous dispersion of the amine under standard conditions of concentration and temperature and at a series of pHs, with plotting of the mobilities against pH to determine the pH of zero mobility. The isoelectric points of a series of fatty amines are given in the following table, in which T and Co represent the mixed alkyl radical derived from hydrogenated tallow acid and coconut oil acid, respectively. From this table it can be seen that those tertiary amines which are members of the above special group have the required isoelectric points, whereas the other amines do not.

Amines of the special group are in general old compounds, and their use has been described for purposes other than as fabric-softening agents. They can be prepared by the processes described in Kirk-Othmer's Encyc-.

lopaedia of Chemical Technology, Volume 2, under Fatty Amines, or in US Patent 3,471,562, which describes the preparation of dioctadecylmethylamine and its use as an intermediate for the preparation of quaternary ammonium compounds which are themselves cationic fabric softeners but do not possess the property of charge reversal. British Patent 1,286,054 and US Patent 3,696,056 describe the use of fatty amines in compositions for controlling foam generated by synthetic detergent compositions and make no mention of fabric softening; the only tertiary amine having the structure given above that is specifically mentioned is a substance designated methyl dihydrogenated tallow amine.

T ₂ NH	7.4	
T ₃ N	8.2	
T2NCH3	8.8	
T2N(CH2)3CH3	9.0	85
CO2NCH3	9.1	
T 2NCH2CH3	9.4	
TN(CH ₃) ₂	10.0	
TNH2	10.9	-
		Or i

The only compositions disclosed containing that amine are ineffective for the purpose of providing fabric-softening, and in order to obtain the practical benefit of the present invention it has been found essential to formulate the fabric-softening amines in such a way as to provide compositions which are different from those described in British Patent 1.286.054 and US Patent 3.696.056.

1,286,054 and US Patent 3,696,056.
According to the present invention, a fabric-softening composition comprises a tertiary amine having an isoelectric point of from 8.3 to 9.8 and of the structure RR¹R²N where R is an alkyl group having from 1 to 6 carbon atoms and R¹ and R² are primary linear alkyl or alkenyl groups having from 10 to 26 carbon atoms, and a diluent, in which the diluent is water or a water-soluble non-surface-active salt; and where the diluent is water the composition is liquid and comprises from 2 to 50% of the amine dispersed in from 98 to 50% of water, by weight of the total amount of the amine and water, and where the diluent is a water-soluble non-surface-active salt, the composition is solid and comprises from 6.3 to 90% of the amine and from 10 to 93.7% of the salt by weight of the composition.

The tertiary amine softeners are preferably those where R has from 1 to 4 carbon atoms, and especially those where R is a methyl group. R will normally be a primary alkyl group. Preferably R¹ and R² each have from 12 to 22, and especially from 16 to 18, carbon atoms. R¹ and R² are linear groups but similar groups with minor amounts of branching insufficient to alter the character of the compounds as softeners are obvious chemical equivalents. Mixtures of amines can be employed, for instance commercial mixtures of compounds in which the alkyl or alkenyl

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groups are derived from the fatty acids of natural fats, as when coconut and hydrogenated tallow acids are converted to their nitriles and these are subsequently hydrogenated and alkylated. In accordance with usage in the fatty amine art, such mixed amines are here referred to as "coco" and "hydrogenated tallow" amines. R¹ and R² can be similarly derived from soyabean oil fatty acid. Preferably R¹ and R² are groups derived from a common source so that in single compounds they are the same group, and in commercial mixtures of compounds they represent the same individual groups in the same proportion.

Examples of suitable amines for use as wash-cycle softeners are:

Methyldidecylamine Methyldilaurylamine Methyldimyristylamine Methyldicetylamine Methyldimargarylamine Methyldistearylamine Methyldiarachidylamine

25 Methyldidocosylamine Methylditetracosylamine Methyldicocoamine Methyldi(hydrogenated tallow)amine

Ethyldilaurylamine
Etyhyldicetylamine
Propyldistearylamine
Butyldistearylamine
Ethyldicocoamine

Ethyldi(hydrogenated tallow)amine
Butyldi(hydrogenated tallow)amine
Methyldicocoamine and methyldi(hydrogenated tallow)amine are particularly suit-

genated tallow)amine are particularly suitable.

The invention provides a process for preparing an aqueous amine dispersion of the invention, in which the amine is dispersed in

water in the presence of an emulsifier, preferably at a temperature at which the amine is in the liquid form. Suitable emulsifiers are listed in Kirk-Othmers' Encyclopaedia of Chemical Technology, Volume 8, page 128-130. The resulting emulsion may on cooling become a suspension due to solidification of the amine. Most dispersions thus

prepared have a short life because they are unstable, and if stored before use are liable to separate into an aqueous phase and a solid amine phase: the successful preparation of dispersions that are stable for many weeks has presented a problem, for it has been found that the use of the standard procedures of emulsion technology for choosing

an emulsifier system having the optimum hydrophiliclipophilic balance (HLB) fails to produce stable emulsions. It has however been discovered that satisfactorily stable dispersions can be prepared by introducing an emulsifier with a hydrophobic group

similar to the hydrophobic groups of the amine; this appears to function by enabling

the formation of amine particles having at their surface the anionic polar head group of the emulsifier carrying a negative charge, thus altering the effective isoelectric point of the amine, while the packing of the hydrophobic group of the emulsifier with the corresponding groups of the amine enables retention of this group within the particles.

Accordingly a liquid composition of the invention preferably contains as emulsifier an anionic surface-active agent having a hydrophobic group that is a primary linear alkyl or alkenyl group having an average chain length within 21/2 carbon atoms of the average chain length of the groups R1 and R2 in the amine. Particularly suitable are emulsifiers having a hydrophobic group distribution that is substantially the same as that of the amine. The anionic head group of the emulsifier can be carboxylate, sulphate, sulphonate, phosphate or phosphonate. Suitable emulsifiers can be chosen from sodium laurate, myristate, palmitate and stearate, disodium hexadecenyl succinate (which has an effective hydrophobic chain length of 16 carbon atoms), sodium N-tallow acyl glutamate (where the "tallow acyl" is a mixture of the acyl groups of tallow fatty acids, and the emulsifier has an effective hydrophobic chain length of 17 carbon atoms), sodium C16_C18 sulphonate (having an effective hydro-phobic chain length of 17 carbon atoms), sodium tallow fatty acid sulphonate (having an effective hydrophobic chain length of 17 carbon atoms) and sodium C16_C20 n-alkyl phosphate (having an effective hydrophobic chain length of 18 carbon atoms). Other emulsifiers which can be used are sodium hexylbenzene sulphonate and sodium octadecylbenzene sulphonate. Instead of sodium salts the salt of any equivalent cation providing water-solubility, for instance lithium, potassium or ammonium, can be used.

Where the amine employed is methyldicocoamine, the emulsfier is preferably sodium laurate, and especially the sodium salt of coconut oil fatty acid. Where the amine is methyldi(hydrogenated tallow)amine the emulsifier is preferably sodium stearate and especially the sodium salt of hydrogenated tallow acid.

Preferably the liquid amine composition contains from 5 to 30%, and especially from 7 to 25%, of the amine by weight of the total amount of amine and water, as such concentrations confer practical advantages in terms of correspondence to existing habits of users of liquid fabric conditioners having regard to the need to dispense a reasonable bulk of liquid in standard amounts and to provide an effective amount of amine in the wash liquor. An effective amount of the emulsfier is chosen from

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within the range of 0.002 to 10% by weight of the dispersion, and generally from 0.1 to 20% by weight of the disperse phase is employed

Preferably the solid amine composition contains at least 6.5, and especially at least 7%, of the amine. The amount of the salt present will usually be less than 75% by

weight of the composition.

As water-soluble non-surface-active salts there are preferably used detergency builder salts. By "detergency builder salt" in this specification is meant a non-surface-active salt that possesses the power of increasing the detergent activity of a detergent-active compound. Suitable detergency builder salts are inorganic builders such as sodium ortho-, pyro-, trimeta- and tripoly- phosphates, sodium carbonate and sodium silicate. Examples of organic builders are salts of organic acids such as sodium citrate, sodium oxydiacetate, sodium carboxymethyloxysuccinate, sodium nitrilotriacetate and sodium ethylenediamine tetraacetate.

Non-surface-active salts other than detergency builders which can be used in solid compositions of the invention are sodium perborate or percarbonate which provide a bleaching function, and sodium

sulphate.

Preferably where the composition contains a detergency builder salt, it also contains an anionic detergent-active compound, especially a non-soap (or synthetic) anionic detergent-active compound. Such a composition is preferably one containing from 6.5 to 35% of amine, from 3 to 60% of anionic detergent-active compound, and from 10 to 75% of detergency builder salt, by weight of

the composition.

Where a composition of the invention contains an anionic detergent-active compound, it can be a water-soluble or waterdispersible alkali metal salt of an organic acid, especially a sodium or potassium salt, or a corresponding ammonium or substituted ammonium salt. Examples of suitable organic acids are alkylbenzene sulphonic acids whose alkyl groups contain from 8 to 20 carbon atoms, for instance linear C10-C15 alkylbenzene sulphonic acids; alkyl and alkenyl sulphonic acids of from 8 to 22 carbon atoms, for instance those known as olefin sulphonates which can be derived by reaction of sulphur trioxide with linear and branched olefins, especially "cracked wax" or "Ziegler" alpha-olefins, or those derived by reaction of alkanes with sulphur dioxide and chlorine and subsequent hydrolysis, or by reaction of olefins with bisulphites; alkyl sulphosuccinates derived by reacting maleic acid esters with bisulphites; alkyl sulphuric acids of from 8 to 22 carbon atoms obtained 65 by reaction of alcohols and sulphur trioxide;

alkylether sulphuric acids obtained by reaction of molar quantities of alcohols of from 6 to 18 carbon atoms with 1 to 15 mols of ethylene oxide or mixtures of ethylene oxide and propylene oxide, and subsequently reacting the condensation product with sulphur trioxide; natural or synthetic aliphatic carboxylic acids of from 10 to 22 carbon atoms, especially the soaps obtained by splitting of triglyceride oils: and N-acylated isethionic and sarcosinic acids, where the acyl groups are those derived from such aliphatic carboxylic acids. Other suitable anionic detergent-active compounds are described in "Surface Active Agents", Volume I by Schwarz and Perry and "Surface Active Agents and Detergents", Volume II by Schwartz, Perry and Berch.

Compositions of the invention can contain other fabric-washing detergent composition ingredients, for instance lather boosters, such as coconut monoethanolamide; lather controllers; chlorine-releasing bleaching agents, for instance trichloroisocyanuric acid and sodium and potassium dichloroisocyanurates; antiredeposition agents, such as sodium carboxymethylcellulose; and perfumes, colourants, fluorescers, corrosion inhibitors, germicides and enzymes.

The invention provides a process for preparing a solid composition of the invention in which the amine is mixed with the nonsurface-active salt and anionic detergentactive compound where present and, if required, the mixture is formed into granules. Preferably the solid composition is prepared as a spray-dried powder. Such a powder can be obtained by preparing an aqueous slurry containing the ingredients of the composition and heating the slurry to 80°C, at which the amine will be liquid and can with adequate mixing be distributed uniformly throughout the slurry, which is then spray-dried to give a powder containing evenly-distributed amine.

For domestic use, a liquid composition of the invention can be added in small quantities sufficient to provide softening, for instance from a bottle or a sealed sachet, for example from 10 to 100 ml, to a fabric load in a washing machine, together with a suitable detergent composition, for instance a washing powder. Solid compositions of the invention without the necessary detergent for a wash are added in the same way, and solid compositions containing detergentactive compound for a wash are used in the same way as normal detergent compositions. In order to get good results the pH of the wash liquor should be at least 0.2 pH units above the isoelectric point of the amine employed. This can be achieved by selection of a suitable amine for a given wash liquor or by adjusting the alkali content of the wash liquor, if required. The pH

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	of the wash liquor is preferably below 9.8 in	Fluorescer 0.2	
	order to avoid too great a buffering effect to	Water 10.0	
	enable the necessary reduction of pH in the	The resulting load had a liquor:cloth ratio	
	rinse liquor to be made without having to	of 7:1 with the wash liquor at pH 9.3. The	
5	lower the pH of natural rinse water: how-	fabrics were washed with a standard prog-	70
_	ever, a wash liquor of high pH can be	ramme in which the load was heated to 95°	•
	diluted with water after washing and before	with drum agitation during 35 min and then	
	wash liquor discharge if it is desired to	drained, leaving about 20% of residual wash	
	reduce the contribution of the residual wash	liquor, and rinsed four times with cold water	
10	liquor to the alkalinity of the first rinse	(20 litres, pH 7.5, giving pH 8.7 in the first	75
10	liquor. For practical purposes the wash	rinse liquor), with spin-drying after the third	
	liquor pH should be above 8.5, so that scope	and fourth rinse. The cloth was then allowed	
	is given for use of an amine whose isoelec-	to dry in the atmosphere. The dried cloth	
	tric point is at least 0.2 units above the pH	had a surface-coating of about 0.002 parts	
15	of the first rinse liquor.	of amine per part by weight of the fabric.	80
	The invention includes a process for the	A similar operation was also carried out	
	softening of fabrics, in which an aqueous	without the amine dispersion. The cloth	
	anionic detergent wash liquor is prepared	pieces were assembled in pairs, one washed	
	incorporating a composition of the inven-	with amine and one without, and directly	
20	tion, the liquor having a pH above the	compared by a test panel of 5 operators who	85
	isoelectric point of the amine, fabric is	alloted an order of softening preference to	
	washed with the wash liquor, and the pH of	each pair. In 77 out of 80 pair comparisons	
	the wash liquor in contact with the washed	the cloth treated with the amine dispersion	
	fabric is then reduced to below the isoelec-	was selected as softer.	20
25	tric point of the amine to attach the amine to		90
	the surface of the fabric. Preferably the bulk	EXAMPLE 2	
	of wash liquor present during the washing	An amine dispersion was prepared as in	
	step is separated from the washed fabric and	Example 1, but using as amine	
	the pH reduction is effected by the addition	methyldi(hydrogenated tallow)amine and as	ne.
30	of rinse water to the fabric containing the	emulsifier sodium stearate.	95
	residual wash liquor.	Soiled laundry (2.7kg) and 16 pieces	
	The invention also includes a softened	(20.3cm square) of cotton towelling was	
	fabric having a surface-coating of from	placed in a top-loading paddle washing	
	0.0005 to 0.01 , and preferably from 0.001	machine, followed by water (68 litres) at	300
35	to 0.005, parts per part by weight of the	50°, the amine dispersion (50ml) and a	100
	fabric, of a tertiary amine of the special	detergent product (100g) having the follow-	
	group defined above having an isoelectric	ing composition in parts by weight:	
	point of 8.3 to 9.8.	Sodium dodecylbenzene sulphonate 6.0	
40	The invention is illustrated by the follow-	Condensate of hardened tallow	105
40	ing Examples, in which all temperatures are	fatty alcohol with 18 molar equivalents of ethylene oxide 4.0	105
	in °C.		
	EXAMPLE 1	22.0	
	To a solution in water (84.85g) of sodium	Sodium tripolyphosphate 33.0 Alkaline sodium silicate 8.0	
15	laurate (0.15g) at 80° was added molten	Sodium sulphate 3.1	110
45	methyldicocoamine (15g) and the mixture	Sodium carboxymethyl cellulose 1.0	
	stirred and subjected to ultrasonic disper-	Sodium perborate 25.0	
	sion until it was homogeneous. Soiled laundry (2.7kg) and 16 pieces	Fluorescer and perfume 0.7	
	(20.3cm square) of clean cotton towelling	Water 11.2	
50	were placed in a front-loading automatic	The resulting load had a liquor : cloth	115
50	drum washing machine, followed by water	ratio of 25:1, with the wash liquor at pH 9.7.	
	(20 litres) at 50°, the amine dispersion	The fabrics were washed with a standard	
	(75ml) and a detergent product (100g) hav-	programme in which the load was heated to	
	ing the following composition in parts by	95° with paddle agitation during 40 min and	
55	weight:	was drained and excess water removed by	120
33	Sodium dodecylbenzene sulphonate 7.5	spinning, leaving about 10% of residual	
	Condensate of tallow fatty acid	wash liquor, and rinsed twice with cold	
	amine with 11 molar equivalents	water (60 litres, pH 7.5, giving pH 8.2 in the	
	of ethylene oxide 3.5	first rinse liquor), with spin-drying after the	
60		final rinse, and the cloth allowed to dry in	125
	Sodium tripolyphosphate 64.9	the atmosphere. The dried cloth had a	
	Alkaline sodium silicate 1.0	surface-coating of about 0.0015 parts of	
	Sodium sulphate 9.5	amine by weight of the fabric.	
	Sodium carboxymethyl cellulose 1.0	The procedure was repeated using 32	4.5.
65	Sodium chloride 1.4	pieces of towelling and no amine dispersion:	130
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genated tallow)amine

	16 were removed and dried and the machine set for one more rinse and spin cycle, and	prepared as in Example 2, except that the amine used was a methyldialkylamine of
	50ml of a commercial rinse conditioner con-	isoelectric point 9.3, each of whose alkyl
	taining 6% by weight of a mixture of	groups was derived from a commercial mix-
5	dimethyldicocoammonium chloride and	ture of C ₂₀ and C ₂₂ straight chain saturated
	dimethyldi(hydrogenated tallow)	fatty acids in which the C20 acid predomi-
	ammonium chloride as softener was added	nated.
	and the 16 remaining pieces dried.	Similar dispersions were prepared for
	The 16 pieces from each treatment were	comparisons using as amine tri(hydroge-
10	arranged in triplets and submitted to a soft-	nated tallow)amine and dimethyl(hydroge-
	ness comparison by a 5-membered test	nated tallow)amine whose isoelectric points
	panel. Each triplet was allotted a score of 1	are 8.2 and 10.0 respectively. The amine
	for the softest piece, 2 for the next softest	dispersion of Example 2 was also used
	and 3 for the harshest. The total softness	(Example 5).
15	scores were as follows.	The washing procedure of Example 3 was
	Cloths treated with	carried out with each emulsion (with wash
	amine dispersion 130	liquid pH 9.7 and first rinse liquor pH 8.2).
	commercial rinse conditioner 113	The cloths were divided into 16 sets of 4
••	detergent only 237	pieces including one washed with each
20	This demonstrates that softening with the	emulsion, and graded by a test panel of 5
	amine dispersion used as a wash cycle sof-	persons for softness, with scores allotted as
	tener was not significantly poorer than that	I for the softest and 4 for the harshest. The combined softness scores obtained
	with the commercial quaternary ammonium	were:
25	salt composition when used as a rinse cycle softener, both showing marked softening.	Cloths treated with dispersions of
25	Softener, both showing marked softening.	methyldi(C20-C22 alkyl)amine
	EXAMPLE 3	(Example 4) 136
	An amine dispersion was prepared as in	methyldi(hydrogenated
	Example 1, but using as amine methyldi-	tallow)amine
30	stearylamine (isoelectric point 8.7) and as	(Example 5) 167
-	emulsifier sodium stearate.	tri(hydrogenated tallow)amine 301
	As detergent there was used a composi-	dimethyl(hydrogenated
	tion having the following ingredients in	tallow)amine 196
	parts by weight:	The scores with the amine dispersions of
35	Sodium dodecylbenzene sulphonate 14.0	examples 4 and 5 demonstrate a softening
	Coconut ethanolamide 2.0	effect in comparison with those with the
	Sodium tripolyphosphate 33.0	other dispersions. The cloths washed with
	Alkaline sodium silicate 6.0	the last dispersion had greasy deposits due
	Sodium sulphate 17.0 Sodium carboxymethylcellulose 0.5	to a precipitate formed from the amine and
40		the anionic detergent used.
	Sodium perborate 20.0 Fluorescer and perfume 0.2	EVAMBLE 4
	Water 7.3	EXAMPLE 6
	In a Tergotometer were placed 4 pieces of	The amine dispersion of Example 2 was
45	clean cotton towelling (20.3cm square),	used as a wash cycle softener as in Example 3, in comparison with a commercial wash
7.7	water (800ml), amine dispersion (2ml) and	cycle softener containing 14% by weight of
	detergent composition (3.2g), giving a	a quarternary imidazoline methyl sulphate,
	liquor: cloth ratio of 20:1, with the wash	with the commercial quaternary ammonium
	liquor at pH 9.7. The cloth was washed for	salt rinse conditioner of Example 2, and also
50	15 min at 80°, the wash liquor discharged,	with the detergent composition of Example
	leaving about 10% as residue, the cloth	3 with no added softener. The procedure
	rinsed twice in cold water (800ml, 24°C, pH	used was similar to that in Example 3 except
	7.5, giving pH 8.2 in the first rinse liquor),	that 1ml of the amine emulsion of Example
	and dried. The dried cloth had a surface-	2 and 1ml of the quaternary imidazoline salt
55	coating of about 0.002 parts of amine by	softener were used as wash cycle softeners,
	weight of the fabric.	and washed were also carried out with the
	The procedure was also carried out using	detergent composition with no softener both
	no amine dispersion. The sets of cloths were	with and without the commercial rinse con-
	tested for softness in pairs as in Example 1.	ditioner of Example 2 (1ml) added to the
60	Every member of the panel found that the cloth treated with amine dispersion was sof-	last rinse liquor. The resulting dried cloths
	ter than that treated with detergent alone.	were panel-tested as before, and the follow-
	tor than that treated with detergent aione.	ing softness scores obtained.
	EXAMPLES 4 and 5	Cloths treated with
	LAMIN LLS 4 and 3	dispersion of methyldi(hydro-

An amine dispersion (Example 4) was

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-	comme	rcial wash cycle softener rcial rinse conditioner	270 142	ing the following ingredients in parts by weight was prepared.	
	deterge		280	Sodium dodecylbenzene sulphonate 14.0 Coconut ethanolamide 2.0	
_		EXAMPLES 7 to 15	;	Sodium tripolyphosphate 33.0	70
5	Amir	e dispersions were pre		Sodium sulphate 6.9	
	Exampl	le 2, but using a range of	amine con-	Alkaline sodium silicate 6.0	
	centrati	ons, namely 5, 10, 20,	25, 30, 35,	Methyldi(hydrogenated	
	40, 45	and 50% of methyldi(hy	drogenated	tallow)amine 10.0	75
0	tallow)	amine by weight of the	dispersion	Sodium carboxymethylcellulose 0.5	15
	and so	dium stearate as emu	isilier. The	1140105001	
		ons were found to be so weeks at ambient tempe		Sodium perborate 20.0 Water 7.2	
		process of Example 3 using		All the ingredients except the amine and the	
5	dispersi	on but with the amine	dispersion	perborate were mixed together with an	80
J	replace	d by 1ml of the 30%	dispersion	additional amount of water (62 parts) to	
	above	(Example 11) was carri	ed out and	provide a slurry, and heated to 80°, the	
		d cotton fabric produced		amine was added in the molten state and the	
		-		mixture thoroughly stirred and spray-dried	0.5
0		EXAMPLES 16 to 2		to give a dry free-flowing powder, with	85
	Amiı	ne dispersions were pre	pared as in	which the perborate was finally mixed.	
	Examp	le 2, but using a range of	of emulsifier	The composition was used to wash cotton pieces in a Tergotometer at 0.4% concent-	
	concen	trations, namely 0.015,	U.43, U./3,	ration as in Example 3, with a wash liquor	
_	1.5 and	13% of sodium stearate persion. The dispersions	were found	pH of 9.7 and a first rinse liquor pH of 8.3.	90
5	the disp	table for at least 6 weeks	s at ambient	The dried cloth had a surface-coating of	, ,
	temper	ature, and that with 0.	45% emul-	about 0.004 parts of amine by weight of the	
	sifier w	as stable for at least 6 n	onths.	fabric. A control wash was also carried out	
	511101			using a similar composition in which the	
0		EXAMPLES 21 to 3	30	amine was replaced by the same amount of	95
	Ami	ne dispersions were pre	pared as in	additional sodium sulphate. Comparison by	
	Examp	le 2, but using a series	of different	a test panel as before was made and the	
	emulsi	fiers, and the period of	stability of	cloth treated with amine was chosen as sof-	
_	_	mulsion at 20° was obse	rved, as fol-	ter in every instance.	10
5	lows.	I. Elaifean	Stability	EXAMPLE 34	10
	Examp		Stability 10 days	A solid fabric-softening composition was	
	21 22	sodium laurate sodium myristate	2 weeks	prepared having the same ingredients as the	
	23	sodium palmitate	at least	detergent product of Example 1 except that	
0	45	Sourain paining	6 weeks	7.5 parts of methyldi(hydrogenated tal-	- 10
•	24	sodium hexadecenyl	at least	low)amine was substituted for the same	
		succinate	10 weeks	amount of sodium sulphate. The composi-	
	25	sodium N-tallow	4 weeks	tion was prepared by mixing together all the	
_		acyl glutamate	4	ingredients except the amine and a portion (23 parts) of the sodium tripolyphosphate	11
5	26	sodium Cia-Cia	4 weeks	together with an additional amount of water	
	27	a-olefin sulphonate	at least	(60 parts) to provide a slurry, heating the	
	27	sodium hexylbenzene sulphonate	10 weeks	mixture to 80°, adding the molten amine	,
	28	sodium octadecylben-	4 weeks	stirring the mixture thoroughly and spray-	
0	20	zene sulphonate		drying to give a dry free-flowing powder	. 11
J	29	sodium tallow fatty	at least	with which the remaining tripolyphosphate	:
	-	acid sulphonate	10 weeks	was mixed.	
	30	sodium C16-C20 n-alkyl	6 weeks.	The composition was used to wash cotton	
		phosphate		pieces in a Tergotometer as in Example 3 a 0.2, 0.4 and 0.6% concentrations, with a	1 12
5				wash liquor pH of 9.3 and a first rinse liquo	. 12
		EXAMPLES 31 and		pH of 8.1, and control washes were also car	
	Stab	Stable amine dispersions are prepared as in Example 1 but using as amine and emul-		ried out using the detergent product o	f
	in Exa	mple I but using as amil	roamine and	Example 1. Comparison by a test panel a	3
	sitter	respectively methyldicood fium salt of coconut oil fa	tty acid- and	before showed that the amine-treated cloth	1 12
0	INC SOC	ldi(hydrogenated tallow	namine and	was softer than the control in every instance	
	the so	dignydrogenated tanow	d tallow fatty	•	
	acid.	mani san or nyorogenate		EXAMPLE 35	
	aciu.	EXAMPLE 33		A composition is prepared as in Example	;
65	·Asc	olid fabric-softening com	position hav-	34, except that the amine employed i	13
		0 -		-	

methyldicocoamine: 100g of the composition is used in a front-loading automatic washing machine in a laundering process similar to that described in Example 1, with a wash liquor pH of 9.3 and a first rinse liquor pH of 8.7, to provide softening of the cotton fabric.

WHAT WE CLAIM IS:--

1. A fabric-softening composition comprising a tertiary amine having an isoelectric point of from 8.3 to 9.8 and of the structure RR'R'N where R is an alkyl group having from 1 to 6 carbon atoms and R' and R' are primary linear alkyl or alkenyl groups having from 10 to 26 carbon atoms, and a diluent, in which the diluent is water or a water-soluble non-surface-active salt; and where the diluent is water the composition is liquid and comprises from 2 to 50% of the amine dispersed in from 98 to 50% of water, by weight of the total amount of amine and water, and where the diluent is a water-soluble non-surface-active salt, the composition is solid and comprises from 6.3 to 90% of the amine and from 10 to 93.7% of the salt by weight of the composition.

2. A composition according to Claim 1, where R in the amine has from 1 to 4 carbon

30 atoms.

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3. A composition according to Claim 2,

where R is a methyl group.

4. A composition according to any preceding claim, where R1 and R2 each have 35 from 12 to 22 carbon atoms.

5. A composition according to Claim 4, where R1 and R2 each have from 16 to 18 carbon atoms.

6. A composition according to Claim 3, where the amine is methyldicocoamine.

7. A composition according to Claim 3, where the amine is methyldi(hydrogenated tallow)amine.

8. A composition according to any pre-45 ceding claim, in which the diluent is water.

9. A composition according to Claim 8, and containing from 5 to 30% of the amine.

10. A composition according to Claim 9, and containing from 7 to 25% of the 50 amine.

A composition according to any one of Claims 8 to 10, and containing as emulsifier an anionic surface-active agent having a hydrophobic group that is a primary linear alkyl or alkenyl group having an average chain length within 2½ carbon atoms of the

average chain length of the groups R1 and R² in the amine.

12. A composition according to Claim 11, where the amine is methyldicocoamine and the emulsifier is sodium laurate.

13. A composition according to Claim 11, where the amine is methyldi(hydrogenated tallow)amine and the emulsifier is sodium stearate.

14. A composition according to Claim 11, in which the emulsifier has a hydrophobic group distribution that is substantially the same as that of the amine.

15. A composition according to Claim 14, where the amine is methyldicocoamine and the emulsifier is the sodium salt of

coconut oil fatty acid.

16. A composition according to Claim 14, where the amine is methyldi(hydrogenated tallow)amine and the emulsifier is the sodium salt of hydrogenated tallow acid.

17. A composition according to any one of Claims 1 to 7, in which the diluent is a water-soluble non-surface-active salt.

18. A composition according to Claim 17, and containing at least 7% of the amine.

19. A composition according to Claim 17 or Claim 18, in which the salt is a detergency builder salt.

20. A composition according to Claim 19, and containing an anionic detergent-

active compound.

21. A composition according to Claim 20, and containing from 6.5 to 35% of the amine, from 3 to 60% of anionic detergent-active compound, and from 10 to 75% of detergency builder salt, by weight of the composition.

22. A composition according to any one of Claims 17 to 21, and containing as salt

sodium tripolyphosphate.

23. A composition according to any one of Claims 17 to 22, and containing as salt sodium perborate or percarbonate.

24. A composition according to any one of Claims 17 to 23, and containing as salt sodium sulphate.

25. A composition according to Claim 1, substantially as described in any one of

the accompanying Examples.

26. A process of preparing a composition according to any preceding claim, in which the amine is dispersed in water in the presence of an emulsifier or is mixed with 110 the water-soluble non-surface-active salt.

27. A process according to Claim 26, substantially as described in any one of the

accompanying Examples.

28. A process for the softening of fabrics, in which an aqueous anionic detergent wash liquor is prepared incorporating a composition according to any one of Claims 1 to 25, the liquor having a pH above the isoelectric point of the amine, fabric is washed with the wash liquor, and the pH of the wash liquor in contact with the washed fabric is then reduced to below the isoelectric point of the amine to attach the amine to the surface of the fabric.

29. A process according to Claim 28, in which the bulk of wash liquor present during the washing step is separated from the washed fabric and the pH reduction is effected by the addition of rinse water to the

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75

80

85

100

120

fabric containing the residual wash liquor.

30. A process according to Claim 29, substantially as described in any one of Examples 1 to 6, 11 and 33 to 35.

31. A softened fabric when obtained by a process according to any one of Claims 28 to 30.

to 30.

32. A softened fabric having a surface-coating of from 0.0005 to 0.01 parts, per part by weight of the fabric, of an amine as defined in Claim 1.

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